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FILE CONTENT: 1840 - 10 Oct 2004 VOL 141 ISS 15

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d que L1

STR

Structure attributes must be viewed using STN Express query preparation.

L3 0 SEA FILE=CASREACT SSS FUL L1 (0 REACTIONS)

=> => file caplus
FILE 'CAPLUS' ENTERED AT 13:22:39 ON 14 OCT 2004
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FILE COVERS 1907 - 14 Oct 2004 VOL 141 ISS 16 FILE LAST UPDATED: 13 Oct 2004 (20041013/ED) This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d que

L4

STR

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L6

3 SEA FILE=REGISTRY SSS FUL L4

L7

2 SEA FILE=CAPLUS L6

=> d 17 1-2 ibib abs hitstr

ANSWER 1 OF 2 CAPLUS COPYRIGHT 2004 ACS on STN L7

ACCESSION NUMBER:

2003:696541 CAPLUS

DOCUMENT NUMBER:

139:230631

TITLE:

Four-step process for the preparation of

3-carboxymethylsulfinyl-7-fluoro-3-methyl-4-quinolone

from flosequinan

INVENTOR (S):

Kwiatkowski, Stefan; Golinski, Miroslaw

PATENT ASSIGNEE(S):

SOURCE:

R.T. Alamo Ventures I, LLC, USA

U.S. Pat. Appl. Publ., 8 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

US 2003166678 A1 20030904 US 2002-281800 20021028 US 6689791 B2 20040210	
US 2003191152 A1 20031009 US 2002-282286 20021028	
PRIORITY APPLN. INFO.: US 2002-360829P P 20020301	
US 2002-360954P P 20020301	
US 2002-361146P P 20020301	
US 2002-361150P P 20020301	
US 2002-403033P P 20020813	

A four-step process for the preparation of 3-carboxymethylsulfinyl-7-fluoro-3-ΑB methyl-4-quinolone from flosequinan is presented.

IT -591781-23-6P - - - - -

RL: BCP (Biochemical process); SPN (Synthetic preparation); BIOL

(Biological study); PREP (Preparation); PROC (Process)

(four-step process for the preparation of 3-carboxymethylsulfinyl-7-fluoro-3-

methyl-4-quinolone from flosequinan)

RN

591781-23-6 CAPLUS
Acetic acid, [(7-fluoro-1,4-dihydro-1-methyl-4-oxo-3-quinolinyl)sulfinyl]-CN (9CI) (CA INDEX NAME)

L7 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2004 ACS on STN

ACCESSION NUMBER:

1996:163892 CAPLUS

DOCUMENT NUMBER:

124:202042

TITLE:

Preparation of 3-aralkylthio-4-hydroxy-2-quinolones

and analogs as NMDA receptor antagonists

INVENTOR(S):

Allgeier, Hans

CODEN: EPXXDW

PATENT ASSIGNEE(S):

Ciba-Geigy A.-G., Switz.

SOURCE:

Eur. Pat. Appl., 27 pp.

DOCUMENT TYPE:

Patent

LANGUAGE:

German

LANGUAGE:

Germ

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT 1	1O.		KIND	DATE	APPLICATION N	ο.	DATE
EP 6854	56		A1	19951206	EP 1995-81034	4	19950523
R:	AT, BE,	CH,	DE, DK	, ES, FR,	GB, GR, IE, IT,	LI, LU,	NL, PT, SE
AU 95203	336		A1	19951214	AU 1995-20336		19950526
CA 2150	545		AA	19951203	CA 1995-21506	45	19950531
FI 95026	550		A	19951203	FI 1995-2650		19950531
NO 9502	L71		A	19951204	NO 1995-2171		19950601
ZA 95045	507		A	19960201	ZA 1995-4507		19950601
CN 11209	538		A	19960417	CN 1995-10617	9	19950601
HU 72608	3		A2	19960528	HU 1995-1598		19950601
US 56333	379		A	19970527	US 1995-45635	8	19950601
JP'08041	1027		A2	19960213	JP 1995-13672	4 、	19950602
BR 95026	547		A	19960423	BR 1995-2647		19950602
PRIORITY APPI	N. INFO.	:			CH 1994-1732		19940602
OTHER SOURCE	(S):		MARPAT	124:20204	12		
GT							

Title compds. [I; R = Z1Z2R5; R1-R4 = H, aliphatic hydrocarbyl, OH, halo, etc.; R5 = Ph, CO2H, alkoxycarbonyl, etc.; Z1 = O, (oxidized) S; Z2 = divalent aliphatic group] were prepared Thus, Me 4-chloroanthranilate was amidated by BrCOCH2Br and the product etherified by Ph(CH2)3SH to give, after cyclization, I [R = (CH2)3Ph, R1 = R3 = R4 = H, R2 = Cl]. I had IC50 of 0.07-1.25µM against 5,7-dichlorokynurenic acid binding at rat cortex and hippocampus membrane preparation in vitro.

=>

IT174455-94-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of 3-aralkylthio-4-hydroxy-2-quinolones and analogs as NMDA receptor antagonists)

RN

174455-94-8 CAPLUS
Acetic acid, [(7-chloro-1,2-dihydro-4-hydroxy-2-oxo-3-quinolinyl)sulfonyl]-CN(9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{C1} & \text{H} & \text{O} \\ & \text{O} & \text{O} \\ & \text{S-CH}_2\text{-CO}_2\text{H} \\ & \text{OH} & \text{O} \end{array}$$